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- (A) Olefin production.
- A process is provided to stabilize and/or reactivate an olefin production catalyst system which comprises contacting an olefin production catalyst system, either before or after use, with an aromatic compound.

BACKGROUND OF THE INVENTION

Field of this invention relates to olefin production and olefin production catalyst system stabilization.

Olfefins, primarfly alpha-olefins, have many uses. In addition to uses as specific chemicals, alpha-olefins are used in polymerization processes either as a monomer or a comonomer to prepare polyolefins, or polymers. Unfortunately, during any catalyzed reaction, such as an olefin production process or a polymerization process, catalyst system activity and productivity can decrease. Furthermore, harsher reaction conditions, such as increased temperatures, can increase catalyst productivity and selectivity, but, conversely, can result in shorter catalyst system file, quicker catalyst system decomposition, and/or catalyst system destabilization. Disposal of spent catalyst system can pose environmental problems and continual addition of hew catalyst system can be cost prohibitive.

SUMMARY OF THE INVENTION

5 Accordingly, it is an object of this invention to improve olefin production catalyst system activity and productivity.

It is another object of this invention to provide a process which will effectively stabilize and/or reactivate an olefin production catalyst system.

It is a further object of this invention to provide a process which will enhance olefin production using an 20 improved, or stabilized, olefin production catalyst system.

It is yet another object of this invention to provide a process to increase the thermostability of an otefin production catalyst system.

In accordance with this invention, a process is provided to stabilize, regenerate and/or reactivate an olefin production catalyst system which comprises contacting an olefin production catalyst system, either 50 before

DETAILED DESCRIPTION OF THE INVENTION

Catalyst Systems

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Catalyst systems useful in accordance with this invention comprise a chromium source, a pyrrotecontaining compound and a metal alkyl, all of which have been contacted and/or reacted in the presence of an unsaturated hydrocarbon. Optionally, these catalyst systems can be supported on an inorpain coxide support. These catalyst systems are especialty useful for the dimerization and trimerization of olelins, such as, for example, eithlyne to 1-hexense.

The chromium source can be one or more organic or inorganic compounds, wherein the chromium oxidation state is from 0 to 6. Generally, the chromium source will have a formula of CrX_m, whorein X can be the same or different and can be any organic or inorganic radical, and n is an integer from 1 to 6. Exemplary organic radicals can have from about 1 to about 20 carbon atoms per radical, and are selected from the group constisting of altry, latcoy, sets, ketnen, and/or amido radicals. The organic radicals can be straight-chained or branched, cyclic or acyclic, aromatic or alightetic, can be made of mixed alightetic, aromatic, and/or oxidos. Exemplary inorganic radicals include, but are not limited to halides, suffets, and/or oxidos.

Specific exemplary chromium(II) compounds include, but are not limited to, chromous bromide, chromous fluoride, chromous chloride, chromium(II) bai(2-eht/hexanasele, chromium(III) assista, chromium-(II) butyrate, chromium(II) neopertanoale, chromium(III) laurate, chromium(III) stearate, chromium(III) ovalate and/or chromium(III) ovanidate)

The pyrrole-containing compound can be any pyrrole-containing compound, or pyrrolide, that will react with a chromium source to form a chromium pyrrolide complex. As used in this disclosure, the term "pyrrole-containing compound" refers to hydrogen pyrrolide, i.e., pyrrole (Ca-lts,N), derivatives of hydrogen pyrrolide, substituted pyrrolides, as well as metal pyrrolide complexes. A "pyrrolide" is defined as a compound comprising a 5-membered, introgen-containing heterocycle, such as for example, pyrrole, derivatives of pyrrole, and mixtures thereof. Broadly, the pyrrole-containing compound can be either affire affired added to the reaction, or generated in-statu.

Generally, the pyrrole-containing compound will have from about 4 to about 20 carbon atoms per molecule. Exemplary pyrrolides are selected from the group consisting of hydrogen pyrrolide (pyrrole), lithium pyrrolide, sedium pyrrolide, potassium pyrrolide, sedium pyrrolide, sedium pyrrolides, because of high reactivity and activity with the other reactants. Examples of substituted pyrrolides; include, but are not limited to, pyrrole-2-carboxylic acid, 2-acetylpyrrole, pyrrole-2-carboxal-dayle, tetrahydrolide, 2.5-dimethyl-pyrole, 2.4-dimethyl-3-ethylpyrrole, 3-acetyl-2-d-dimethyl-pyrole, 2-dimethyl-3-dimethyl-2-pyrolecarboxylate, and midures thereof. When the pyrrole-containing compound contains chromium, the resultant chromium compound can be called a chromium pyrolide.

The most preferred pyrtole-containing compounds used in a trimerization catalyst system are selected from the group consisting of hydrogen pyrrolide, i.e., pyrrole (CsI+SN), 2,5-dimetylipyrrole and/or chromium pyrrolides because of enhanced trimerization activity. Optionally, for ease of use, a chromium pyrrolide can provide both the chromium source and the pyrrole-containing compound. As used in this discbsure, when a chromium pyrrolide is used to form a catalyst system, a chromium pyrrolide is considered to provide both the chromium source and the pyrrole-containing compound. While all pyrrole-containing compounds can produce catalyst systems with high activity and productivity, use of pyrrole and/or 2,5-dimethylpyrrole can produce a catalyst systems with enhanced activity and selectivity to a desired product.

The metal alkyl can bo any heteroleptic or fromcloptic metal alkyl compound. One or more metal alkyls can be used. The alkyl ligand(s) on the metal can be alighetic antifor aromatic. Preferably, the alkyl ligand(s) are any saturated or unsaturated alighatic radical. The metal alkyl can have any number of carbon atoms. However, due to commercial availability and ease of use, the metal alkyl will usually comprise less as than about 70 carbon atoms per metal alkyl milecule and preferably less than about 20 carbon atoms per metal alkyls include, but are not limited to, alkylatumimum compounds, alkylapono compounds alkyls and safty alkylatumimum compounds, alkylapon compounds and/or alkyl filthium compounds. Exemplary metal alkyls include, but are not limited to, n-butyl lithium, butylifithium, dethylmagnesium, diethylrianc, triethylatuminum, riiteobylutaminum, and mbutres thereof.

Preferably, the metal alkyl is selected from the group consisting of non-hydrolyzod, i.e., not precontacted with water, alkylatuminum compounds, derivatives of alkylatuminum compounds, habgenated alkylatuminum compounds, and mixtures thereof for improved product selectivity, as well as improved catalyst system reactivity, activity, anctior productivity. The use of hydrolyzed metal alkyls can result is decreased offerin, io, liquids, production and increased polymer, is, solids, production.

Most preferably, the metal alkyl is a non-hydrolyzed alkylatiminum compound, expressed by the general formulee Alfs, Alfs,X, Alfs,Q, Alfs,QR, Alfs,QR, and/or Al₅Rs,S, wherein R is an alkyl group and X is a halfsgen atom. Exemplary compounds include, but are not limited to, briethylatiminum, tripropylatiminum, tripropylatiminum, diethylatiminum chloride, diethylatiminum boronide, diethylatiminum periodide, ethylatiminum diethoride, ethylatiminum sesquichioride, and mixtures thereof for best catalyst system activity and product selectivity. The most preferred alkylatiminum compound is tribrylatiminum, for bost requettis in catalyst system activity and product selectivity.

Usually, contacting and/or reacting of the chromium source, pyrrole-containing compound and a metal alky is done in an unsaturated hydrocarbon. The unsaturated hydrocarbon can be any aromatic or allphatic hydrocarbon, in a gas, liquid or solid state. Preferably, to effect through contacting of the chromium so source, pyrrole-containing compound, and metal alkyl, the unsaturated hydrocarbon will be in a liquid state. The unsaturated hydrocarbon can have any number of carbon stems per molecule. Usually, the unsaturated hydrocarbon will comprise less than about 70 carbon atoms per molecule, and preferably, less than about 20 carbon atoms per molecule, and preferably, less than about 20 carbon atoms per molecule, and preferably, less than about 30 carbon atoms per molecule, and preferably, less than about 30 carbon atoms per molecule, and preferably, less than about 30 carbon atoms per molecule, and preferably, less than about 30 carbon atoms per molecule, and preferably, less than about 30 carbon atoms per molecule, and preferably, less than about 30 carbon atoms per molecule, and preferably, less than about 30 carbon atoms per molecule, and preferably, less than about 30 carbon atoms per molecule, and preferably, less than about 30 carbon atoms per molecule, and preferably, less than about 30 carbon atoms per molecule, such a service and perferably, less than about 30 carbon atoms per molecule, such and perferably, less than about 30 carbon atoms perferably less than 30 carbon

improve catalyst system stability, as well as produce a highly active and selective catalyst system. The most preferred unsaturated aromatic hydrocarbon is toluene.

It should be recognized, however, that the reaction mixture comprising a chromium source, pyrrotecontaining compound, metal alkyl and unsaturated hydrocarbon can contain additional components which 5 do not adversely affect and can enhance the resultant catalyst system, such as, for example, halides.

Catalyst System Stabilization

Even though the catalyst system preferably can be produced in the presence of an unsaturated hydrocarbon, in accordance with this invention, additional, beneficial catalyst system stabilization can be achieved, by contacting the catalyst system with an aromatic compound.

The contacting of the catalyst system and aromatic compound can occur at anytime prior to contacting the catalyst system with one of the reactants, such as, for example, an olefin like ethylene, and prior to the introduction of heat to the catalyst system. Therefor, the aromatic compound and the catalyst system 15 preferably can be precontacted outside of the reactor, or contacted in-situ in the reactor. Contacting of the aromatic compound and catalyst system can occur under any conditions sufficient to stabilize the catalyst system in the presence of heat. Generally, contacting temperatures can be within a range of about -50° to about 70 °C, preferably within a range of about 10 to about 70 °C. Most preferably, for ease of use and safety, contacting can occur at a temperature within a range of 20 to 30 °C. Generally, contacting times 20 will be less than about 1 hour, and preferably for a time within a range of 0.01 seconds to about 10 minutes. Most preferably, the contacting time can be within a range of 0.1 seconds to 30 seconds. Additional contact times do not improve catalyst stability, and shorter contact times can be insufficient to allow complete contacting of the aromatic compound and catalyst system and, therefor, insufficient stabilization of the catalyst system. Any pressure which allows thorough contacting of the aromatic compound and catalyst 25 system can be used. Most preferably, any pressure which can maintain the aromatic compound and catalyst system in liquid form, in order to ensure a thorough contacting, can be used. Most preferably, the contacting is done under a dry, inert atmosphere, so as to not alter the catalyst system.

Any aromatic compound can be used which can stabilize the catalyst system in the presence of heat. Generally, the aromatic compound will have greater than or equal to about 6 canbon atoms per molecule and can have any substitutents attached to the organic ring compound. Preferably, the aromatic compound will have from six to 50 carbon atoms per molecule in order to enable a thorough contacting of the aromatic compound with the catalyst system. The aromatic compound can act as its own solvent or can be soluble in another solvent or carrier. Aromatic compounds with 6 to 50 carbon atoms per molecule are generally more soluble in the reactor modulum at reaction conditions and therefore are sally maintained as a facility.

36 Generally, the amount of aromatic compound added to the reactor can be up to about 15 weight persent, based on the amount of solvent in the reactor, and preferably within a range of about 0.1 to about 10 weight persont. Most preferably, the amount of aromatic compound contacted with the catalyst system is writhin a range of 0.5 to 5 weight persont, based on the volume of solvent in the reactor. To much aromatic compound can inhibit catalyst system activity and insufficient aromatic compound cannot act as a stabilizer to the catalyst system. Expressed in different terms, the moles of aromatic compound per mole of active chromium compound in the catalyst system activity and the professibly, and professibly an amount within a range of 400 to 1,000 can be used for the reasons given above.

45 Reactants

Trimerization, as used in this disclosure, is defined as the combination of any two, three, or more olelins, wherein the number of olelin, Le., carbon-carbon double bonds is reduced by two. Reactants applicable for use in the trimerization process of this invention are olelinic compounds which can a self-sect, i.e., trimerize, to give useful products such as, for example, the self reaction of ethylene can give 1-hexene and the self-reaction of 1,3-butsdiene can give 1-decene and/or 1-decine compounds which can react with other olefinic compounds, i.e., co-trimerize, to give useful products such as, for example, co-trimerization of theylene puls havene can give 1-decene and/or 1-tetradecene, co-trimerization of ethylene puls havene can give 1-decene and/or 1-decreased and/or 1-decosene. For example, the number of olefin bonds in the combination of three ethylene units is reduced by two, to no elief bond, in 1-flexene. In another example, the number of olefin bonds in the combination of two 1,3-butsdiene units, is reduced by two, to two olefin bonds in the combination of two 1,3-butsdiene units, is reduced by two, to two olefin bonds in the combination of two 1,3-butsdiene units, is reduced by two, to two olefin bonds in the combination of two 1,3-butsdiene units, is reduced by two, to two olefin bonds in the olefiniane. As used herein, the term "trimerization" is intended to include dimerization of indellens.

well as "co-trimerization", both as delined above.

Suitable trimerizable olefin compounds are those compounds having from about 2 to about 30 carbon atoms per molecule and having at teast one olefinic double bond. Exemptary mono-1-olefin compounds include, but are not limited to acyclic and cyclic olefins such as, for exemple, ethylene, propylene, 1-5 buteria, 2-buteria, isobutylene, 1-periene, 2-periene, 1-hoxono, 2-hexene, 3-hexene, 1-heptene, 2-heptene, 3-heptene, as and mixtures of any two or more thereof. Exemplary diolefin compounds include, but are not limited to, 1.3-butadiene, 1.4-pentadiene, and 1.5-hexadiene. It branched and/or cyclic olefins are used as reactains, while not withing to be bound by theory, it is bolieved that steric hindrance could hinder the trimerization process. Therefore, the branched and/or or cyclic olefins are prefable than the carbon-carbon double bond.

Catalyst systems produced in accordance with this invention preferably are employed as trimerization catalyst systems.

Reaction Conditions

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The reaction products, i.e., oletin trimers as defined in this specification, can be prepared from the catalyst systems of this invention by solution reaction, slurry reaction, and/or gas phase reaction techniques using convolutional equipment and contacting processes. Contacting of the momenter or monomers with a catalyst system can be effected by any manner known in the art. One convenient method is to suspend the 20 catalyst system in an organic medium and to agilate the mixture to maintain the catalyst system in solution throughout the trimerization process. Other known consistion methods can also be employed.

Reaction temperatures and pressures can be any temperature and pressure which can trimerize the ofelin reactants. Generally, reaction temperatures are within a range of about 0° to about 20° C.

Proferably, reaction temperatures within a range of about 60° to about 20° C and most preferably, within a srange of 80° to 150° C are employed. Generally, reaction pressures are within a range of about atmospheric to about 250° psig. Preferably, reaction pressures within a range of about atmospheric to about 250° psig, Preferably, reaction pressures within a range of about atmospheric to about 250° psig and most preferably, within a range of 30° to 70° psig are employed.

Too low of a reaction temperature can produce too much undesirable insoluble product, such as, for example, polymer, and too high of a temperature can cause decomposition of the catalyst system and reaction products. Too low of a reaction pressure can result in low catalyst system activity.

Optionally, hydrogen can be added to the reactor to accelerate the reaction and/or increase catalyst system activity.

Catalyst systems of this invention are particularly suitable for use in trimortization processes. The sturry process is operatily carried out in an inert dilenst (medium), such as a perfittin, cycloparatin, or aromatic shudocarbon. Exemplary reactor diluents include, but are not limited to, isobutane and cyclohexane. Isobutane can be used to improve process compatibility with other known orderin production processes. However, a homogenous trimortization catalyst system is more soluble in cyclohexane. When the reactant is predominately eithlyen, a temperature in the range of about 0° to about 300° C generally can be used. Preferably, when the reactant is predominately eithlyene, a temperature in the range of about 60° to about 110° C is employed.

Products

The olefinic products of this invention have established utility in a wide variety of applications, such as, for example, as monomers for use in the preparation of homopolymers, copolymers, and/or terpolymers.

The further understanding of the present invention and its advantages will be provided by reference to the following examples.

50 EXAMPLES

Example 1

An exemplary classlyst system was prepared under an inert almosphere (nitrogen) using chromium(ill) 2-ethythexanoste (21.3 mmol C7), 25-dimethylpyrrole (63.8 mmol), ethylaluminum dichloride (85.1 mmol) and triethylaluminum (319 mmol) as follows: Chromium (ill) 2-ethylhoxanoste was dissolved in 100 mL anhydrous toluene and 2.5-dimethylpyrrole added to the resulting dark green solution. In a separate container, ethylaluminum dichloride and triethylaluminum vere mixed together. Then, the aluminum alkyl

solution was poured slowly into the chromium/pyrole solution. The resulting dark yellow-brown solution was stirred for 5 minutes and then the solvent removed in vacuo. The remaining oily liquid was diluted to 150 mt. with cyclohexane and allowed to set overright, under nitrogen, at room temperature and pressure. The solution then was filtered to remove a black precipitate from the filtrate, which contained the catalyst 5 system, and was diluted to a volume of 250 mt. using cyclohexane.

The catalyst system solution and solvent (dodecane) were placed in a 1-liter flask equipped with a nitrogen inlet and a stir bar, under an linert nitrogen atmosphere, and the reactor was brought to the desired temporature.

Ethylene was added until the reactor pressure reached 700 psig; and then ethylene was fed on demand for a specific time period. Ethylene flow was then stopped and the reactor was stowly vonted through a cooled, copper coil. Condensed liquids in the coil and reactor were collected and analyzed. A slight nitrogen purgo was used to aid in removal of volatile product(s) once the reactor pressure had dropped to one atmosphere.

The reactor was then resealed and ethylene was added again until the pressure reached 700 psig; ethylene was fed on demand. The reactor was vented again, as described earlier. Reactor resealing, ethylene pressurization, reactor venting, and sample collection were repeated two or three more times.

Run 101 had 10 mg chromium (Cr) charged and Runs 102-104 had 5 mg Cr charged as part of the

The results of four Funs wherein no additional aromatic compounds were added to stabilize the catalyst system are given below in Table 1. The data in Tables 1 and 2 are a summation, or running total, for all sample fractions collected. The column headors in the Tables, such as Gar and Gar, stand for all of one type of oldrin collected, such as all butiones and all hexenes, unless otherwise indicated. The balance of products collected, to total flow welght percent, were oldring with more than 10 cation atoms per molecule.

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		C	2.9	5.7	10.7	35.7	8.0	9.0	8.0	1.0	19.0	6.0	1.8	6.3	3.2	23.8
5																
			0.2	8.0	1.4	2.9	0.2	0.2	0.3	4.0	1.7	0.1	0.3	9.0	9.0	1.3
10		1-c*,	99.2	99.5	99.2	99.3	7.66	7.66	7.66	7.66	99.7	99.2	99.2	99.3	99.3	4.66
15		C.", Total	96.0	92.8	87.1	60.5	98.7	98.9	98.7	98.3	78.4	98.2	97.1	94.4	95.5	73.9
20		C	0.03	0.01	0.02	<0.01	<0.01	0.01	0.02	0.02	<0.01	0.04	0.03	0.02	0.03	<0.01
26	Table 1	Total Polymer Collect- ed, 8		,	,	1.26	,	•	ı	,	2.47	,	,			1.19
30		Total Olefins Collect- ed. &	226.6	268.6	290.1	295.5	95.0	137.1	177.9	210.8	265.1	39.4	72.4	87.9	98.9	129.2
35		Product- ivity g olefins/ g Cr/hr	22,700	13,400	9,700	7,400	38,000	27,400	17,800	14,100		15,800	14,500	8,800	009'9	
40		Total Time,	09	120	180	240	30	9	120	180	resctor liquid	30	09	120	180	reactor
45		Reactor Temp, °C	06	06	06	06	06	06	06	06		115	115	115	115	115
50		Run(a)	101-A	101-B	101-C	101-D	102-A	102-B	102-C	102-D	102-E	103-A	103-B	103-C	103-D	103-E

Run (a)	Reactor Temp, C	Total Time,	Product- ivity g olefins/ g Cr/hr	Total Olefins Collect-	Total Polymer Collect- ed, g	, kg * \$3 * \$3	C.", Total	1-C ⁶ ,	* ** ***	C 11 ,
¥	140	30	7,500			0.34	98.4	86.66	0.2	0.3
щ	140	09	8,300		•	0.15	98.2	99.3	0.2	9.0
104-C	140	120	000'9			0.2	97.9	99.2	0.3	6.0
Д	140	180	5,600		ı	0.1	7.76	1.66	0.3	1.0
104-E	140	reactor			3.20	<0.01	82.0	0.66	1.3	15.9

Weight percent of $1-C_0^{\pm}$ (1-hexene) is based on the total weight of all hexenes collected. Run 101 bad 10.0 mg catalyst system charged; Runs 102-104 had 5.0 mg catalyst system charged. æ

The data in Table 1 show that as reactor temperature is increased, i.e., from 90 °C to 140 °C, catalyst system activity decreases. The date in Table 1 also show that catalyst system activity decreases over time at constant reactor conditions.

Example 2

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Runs 201-204 used identical catalyst systems, as described in Runs 101-104, and olefin production reaction conditions similar to those in Table 1. However, an aromatic compound added to the reactor with the catalyst system in Runs 201-204. Reactor temperature in Runs 201-204 was 140°C and 5 mg of Gr as part of the catalyst system were charged. In Runs 201 and 202, 15 mL of tokene were action to the reactor along with the olefin production catalyst system and in Runs 203 and 204, 15 mL of the 5th distilled n-butylbenzane were added to the reactor along with the catalyst system. The results of Runs 201-204 are given below in Table 2.

		C. S.	8.0	1.2	1.9	3.4	30.3	9.0	1.2	1.6	2.3	25.6	1.0	1.9	2.5	4.5	39.3
5		TX.	0.1	0.2	0.3	9.0	1.7	0.09	0.2	6.0	7.0	1.4	0.1	0.3	5.0	6.0	2.1
10		1-C**,	89.3	66.3	99.3	86.3	99.3	4.66	4.66	4.66	4.66	7.66	99.1	0.66	0.66	0.66	0.66
15		C. Total,	98.3	8.76	97.0	95.2	66.2	98.6	97.9	4.76	9.96	71.9	7.76	9.96	95.8	93.5	8.44
20		ر د د د د د د د د د د د د د د د د د د د	0.08	0.1	0.1	0.06	<0.01	0.1	0.07	0.1	90.0	<0.01	0.2	0.2	0.2	0.1	<0.01
25	Table 2	Total Polymer Collected, 8	,				0.83	1				0.87					2.38
30		Total Olefins Collected, g	62.9	116.6	154.1	182.9	214.4	59.2	0.96	130.7	152.4	175.6	84.0	163.9	226.6	269.7	307.2
35		Productivity, g olefins/ g Cr/hr	26,400	23,300	15,400	12,200	ı	23,700	19,200	13,100	10,200		33,600	32,800	22,800	18,000	
40		Total Time, mins	30	09	120	180	reactor liquid	30	09	120	180	reactor	30	09	120	180	reactor
45		Run	201-A	201-B	201-C	201-D	201-E	202-A	202-B	202-C	202-D	202-E	203-A	203-B	203-C	203-D	203-E
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5		ε υ Έτι Έτι	1.0	1.6	2.1	3.1	28.5	
10		C. E.	0.2	0.3	7.0	7.0	2.1	
15		1-C**,	99.1	99.1	0.66	0.66	98.7	
10		C. Total,	8.76	0.76	96.2	6.46	59.3	
20		C	0.2	0.2	0.3	0.3	<0.01	
25	Table 2, Cont.	Total Polymer Collected, g		•		•	0.37	ad on the
30 35	티	Total Total Olafins Polymer Collected, g Collected, g	64.7	126.3	185.1	239.8	303.2	exene) is base collected.
46		Productivity, g olefins/ g Cr/hr	25,900	25,300	18,000	15,700		Weight percent of $1-C_e^{\alpha}$ (1-hexene) is based on the total weight of all hexenes collected.
45		Total Time, mins	30	09	120	180	reactor	ight percent tal weight o
so		Run	204-A	204-B	204-C	204-D	204-E	(a) We

As can be seen from the data in Table 2, the addition of an aromatic compound to the catalyst system 55 significantly increases catalyst system activity and total grams of olefins collected, without sacrificing selectivity to 1-hexene.

While this invention has been described in detail for the purpose of illustration, it is not to be construed as limited thereby but is intended to cover all changes and modifications within the spirit and scope thereof.

Claims

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- A process to stabilize an olefin production catalyst system comprising contacting a catalyst system comprising a chromium source, a pyrrole-containing compound and a metal alkyl with an aromatic compound under conditions sufficient to improve the catalyst system thermal stability or activity.
- 3. The process of claim 1 or 2, wherein said metal alkyl is a non-hydrolyzed motal alkyl and is selected from alkyl aluminum compounds, alkyl bronn compounds, alkyl magnesium compounds, alkyl zinc compounds, alkyl lithium compounds, and mixtures of two or more thereof, preferably wherein said non-hydrolyzed metal alkyl is an alkyl aluminum compound, most preferably triethyl aluminum.
 - 4. The process of any of claims 1 to 3, wherein said pyrrole-containing compound is selected from pyrrole, derivatives of pyrrole, alkali metal pyrrolides, saits of alkali metal pyrrolides, and mixtures thereof, particularly selected from hydrogen pyrrolide, 25-dimethylpyrrole, and mixtures thereof.
 - The process of any of the preceding claims, wherein said catalyst system further comprises a hallde source.
- The process of any of the preceding claims, wherein said aromatic hydrocarbon has less than 70 carbon atoms per molecule.
 - A process to produce olefins comprising contacting one or more olefins with a catalyst system obtained by the process of any of the preceding claims.
 - The process of claim 7, wherein said clefin has from 2 to 30 carbon atoms per molecule, preferably wherein said clefin is ethylene.
- A process to trimerize ethylene to 1-hexene comprising contacting ethylene with a catalyst system obtained by the process of any of claims 1 to 8.
 - 10. The process of any of claims 7 to 9, wherein said contacting is carried out at a temperature within a range of 0 to 250 °C and a pressure within a range of atmospheric to 17.6 MPa.